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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

	or agent's file reference 03	FOR FURTHER ACTION	See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)
	ternational application No. CT/MX 03/00027 International filling date (day/month/year) 14.03.2003 Priority date (day/month/year) 14.03.2003		nth/year) Priority date (day/month/year)
A61F2/40	al Patent Classification (IPC) o	both national classification and IPC	<u> </u>
Applicant FERREY	RO IRIGOYEN, ROQUE	HUMBERTO	
1. This Auth	international preliminary ex ority and is transmitted to th	amination report has been prepar le applicant according to Article 3	red by this International Preliminary Examining 36.
2. This	REPORT consists of a total	of 5 sheets, including this cover	·sheet.
×	This report is also accompa been amended and are the (see Rule 70.16 and Section	anied by ANNEXES, i.e. sheets o basis for this report and/or sheet on 607 of the Administrative Instru	of the description, claims and/or drawings which have ts containing rectifications made before this Authority
Thes	annexes consist of a total		icuons under the PC+).
3. This r	eport contains indications r	elating to the following items:	
	_	riating to the following items:	
- Jacob of the opinion		•	
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101	Non-establishment of	opinion with regard to novelty, inv	ventive step and industrial applicability
III I	Non-establishment of Lack of unity of invent Reasoned statement	ion Inder Rule 66 2/21/ii) with regard	ventive step and industrial applicability to novelty, inventive step or industrial applicability;
III II IV E V E	Non-establishment of Lack of unity of invent Reasoned statement	ion Inder Rule 66.2(a)(II) with regard ons supporting such statement	
VI E	Non-establishment of Lack of unity of invent Reasoned statement to citations and explanati Certain documents citations defects in the i	ion Inder Rule 66.2(a)(ii) with regard Ions supporting such statement Indeed In	
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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/MX 03/00027

 Basis o 	f the report
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 With regard to the elements of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)):

	0	escription, Pages			
	1	-17	filed with the demand		
	C	laims, Numbers			
	1	-5	received on 16.03.2005 with letter of 16.03.2005		
	D	rawings, Sheets			
	1/7-7/7		filed with the demand		
2	. W la	lith regard to the lang nguage in which the i	guage , all the elements marked above were available or furnished to this Authority in the international application was filed, unless otherwise indicated under this item.		
	Tł	iese elements were a	available or furnished to this Authority in the following language: , which is:		
			ranslation furnished for the purposes of the international search (under Rule 23.1(b)).		
		the language of pu	blication of the international application (under Rule 48.3(b)).		
		the language of a t Rule 55.2 and/or 5!	ranslation furnished for the purposes of intermediately and the contract of th		
3.	Wi inte	th regard to any nuc ernational preliminary	leotide and/or amino acid sequence disclosed in the international application, the examination was carried out on the basis of the sequence listing:		
			ernational application in written form.		
			he international application in computer readable form.		
		furnished subseque	ently to this Authority in written form.		
			ently to this Authority in computer readable form.		
		The statement that	the subsequently furnished written sequence listing does not go beyond the disclosure application as filed has been furnished.		
		The statement that the information recorded in computer readable form is identical to the written seque listing has been furnished.			
4.	The	amendments have a	resulted in the cancellation of:		
		the description,	pages:		
ł		the claims,	Nos.:		
ļ		the drawings,	sheets:		

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/MX 03/00027

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į	5. 🏻	This report has been estab been considered to go bey	lished a	ed as if (some of) the amendments had not been made, since they have I the disclosure as filed (Rule 70.2(c)).			
		(Any replacement sheet co report.)	ntaining	such amer	dments must be referred :	to under item 1 and annexed to	this
ε	B. Ad	lditional observations, if neces	ssary:				
ı	II. No	n-establishment of opinion	with re	agard to no	valty invantive etch and		
		e questions whether the clain				• • •	
	ob	vious), or to be industrially ap	plicable	have not b	een examined in respect c	an inventive step (to be non- of:	
		the entire international appl	ication,				
	\boxtimes	claims Nos. 5					
		because:					
 the said international application, or the said claims Nos. relate to the following subject matter not require an international preliminary examination (specify): the description, claims or drawings (indicate particular elements below) or said claims Nos. are that no meaningful opinion could be formed (specify): 					aims Nos. relate to the foll ation (specify):	owing subject matter which doe) S
					r said claims Nos. are so uncle	ar	
		the claims, or said claims No could be formed.	os. are	so inadequa	itely supported by the des	cription that no meaningful opin	ilon
	\boxtimes	no international search repo	rt has b	een establi	shed for the said claims N	os. 5	
A meaningful international preliminary examination cannot be carried out due to the failure of the nucle or amino acid sequence listing to comply with the standard provided for in Annex C of the Administrations:						to the failure of the nucleotide anex C of the Administrative	ind/
		the written form has not bee	n furnis	hed or does	not comply with the Stand	dard.	
		the computer readable form	has not	been furnis	hed or does not comply w	ith the Standard.	
V.	Rea cita	ssoned statement under Art tions and explanations sup	icle 35 portinç	(2) with reg g such state	ard to novelty, inventive	step or industrial applicabili	ty;
1.	Stat	ement					
	Nov	elty (N)	Yes: No:	Claims Claims	1-4		
	Inve	entive step (IS)	Yes: No:	Claims Claims	1-4		
	Indu	strial applicability (IA)	Yes: No:	Claims Claims	1-4		
2.	Citat	tions and explanations					
	see	separate sheet					

Form PCT/PEA/409 (January 2004)

INTERNATIONAL PRELIMINARY IT EXAMINATION REPORT - SEPARATE SHEET

International application No. PCT/MX 03/00027

Re Item III

Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

Independent method claim 5 corresponds to independent method claim 6 as filed with the demand. Last said claim, however, was not searched.

Re Item V

Reasoned statement with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- (1) The invention refers to syringes for injection of bone cement.
- (2) The problem is to avoid the disadvantages occurring in prior art devices, such as mechanical deformation by high injection forces of insulin syringes used for said purpose, the need for syringe exchange, exposition of the surgeon's hand to radiation.
- (3) The prior art according to D1 (EP-A-235 905) is a system for remote actuation of an insulin syringe comprising an injection syringe, a pressure exerting body, a hydraulic transmission tube and an manual impulsion and fluid transmission syringe.
- (4) The solution according to the claims is an injection syringe in the form of a commercially available 3 ml hypodermic syringe, a hydraulic tube of 1.0 to 1.5 m length for pressure transmission and a pressure exerting body having a diameter larger than the diameter of the manual impulsion and fluid transmission syringe.
- (5) Provision of said features involves inventive step, since the prior art system is for remote insulin injection for patients suffering from diabetic neuropathy. There is no hint towards adaption of said system to the purpose of cement injection by provision of a 3 ml hypodermic syringe, since injection of insulin according to D1 must be done with an insulin syringe. Further, from D1, it is not made obvious to provide a hydraulic tube for pressure transmission, the tube having a length of 1.0 to 1.5 m, since the purpose of the tube according to D1 is to help in avoiding shifting of the needle by a trembling hand. Finally, in the device according to D1, there is no need for the particular relationship of the diameters, since the injection force for insulin is low.
- (5.1) According to WO-A-9728835, there is provided a system for injection of minor amounts of medicine, in which system the injection syringe is of less size than the actuation syringe. Further, pressure in said system is limited.

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18 CLAIMS

- 1. Hydraulic device for injection of bone cement in percutaneous vertebroplasty, that comprise four main parts, namely: injecting syringe, pressure exerting body, hydraulic transmission tube, an manual impulsion and fluid control syringe; the injection syringe is a commercially available disposable 3 ml hypodermic syringe placed next to the patient; the hydraulic tube for pressure transmission, of 1.0 m to 1.5 m length, placed between the injection syringe and the pressure exerting body; the manual impulsion syringe placed after the hydraulic tube and near the operator, characterized by the pressure exerting body consist of hollow cylindrical body in the form of inverted syringe of larger diameter with an adapted ending like an open bolster with the largest external diameter and two diametrical opposed cuts like oval entry, also in the other end one tip of reduced diameter; an peripheral groove in the internal wall of such bolster, couples tightly the wings of injection syringe in a revolved way; such pressure exerting body has a movable piston on axial direction to define two chambers, namely, internal and external.
- 2.- Hydraulic device of injection of bone cement according to the claim 1, characterized by the cylindrical hollow of pressure exerting body (1), in form of an inverted positioned syringe that renders mechanical advantage to the force exercised in the manual syringe, it has a larger diameter and consists of a joining bolster with internal peripheral groove where are coupled the wings of injecting 3 ml syringe; a body cylindrical lengthened hole of 10 ml of volume that contains a first free camera where the plunger (c) of the injection syringe lodge inside the cylinder until coupled with the moving internal piston (4), and a second internal

camera (5) occupied by a hydraulic fluid, this cameras are separated by such piston (4) surrounded by a rubber cap that seals the internal wall of the body of pressure and avoids leakage of the hydraulic fluid; a final end tip of reduced diameter that is connected in a hermetic way to the hydraulic tube (7).

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3.- Hydraulic device of injection of bone cement according to the claim 2, characterized by the bolster is adapted to receive in a first predetermined position of an oval entry (70) the wings of the injection syringe, and in second position by a 90° turn in a peripheral groove (90), placed in a tight way.

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4.- Hydraulic device of injection of bone cement according to the claim 1, characterized by the manual syringe (8) is a lengthened syringe that has a smaller diameter than the pressure exerting body in a 2/1, 3/1, 4/1 ratio, it is connected in continuation, far from the application point by a hydraulic tube.

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5.- A method of operating the device for injection of bone cement that comprises:

to insert a bone biopsy needle in the place where the bone cement is to be delivered.

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to connect the injecting syringe, loaded with the cement, in continuation of the needle;

to couple in a revolved way, the injecting syringe in the internal peripheral groove of the bolster of the pressure exerting body;

to exert pressure on the plunger of the injecting syringe by means of the force exerted in the plunger of the manual syringe placed in the other end of the

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hydraulic tube, this way, the content of the injecting syringe is injected in the patient's vertebral body;

to retract the plunger of the manual syringe to withdraw the internal piston of the body of pressure in position to receive a new loaded cartridge of bone cement;

to uncouple the injecting syringe from the bolster of the body of pressure; to disconnect the empty syringe from the needle placed in the patient's body;

to couple the new cartridge of bone cement (injecting syringe) in the
needle and bolster of the body of pressure, and repeat the previous steps to
place another new cartridge of bone cement, until completing the filling of the
vertebral body.

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